

A34759 (Sheet 1 of 10)

A34759 (Sheet 2 of 10)

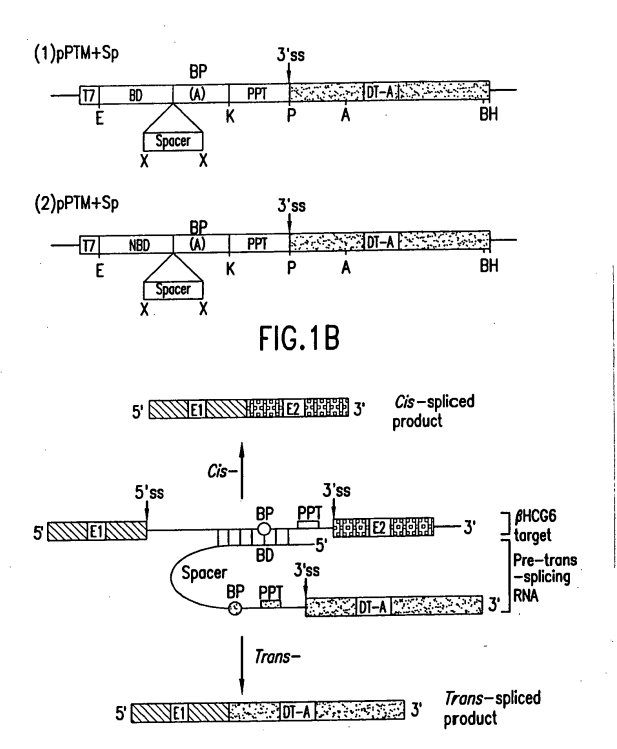


FIG.1C

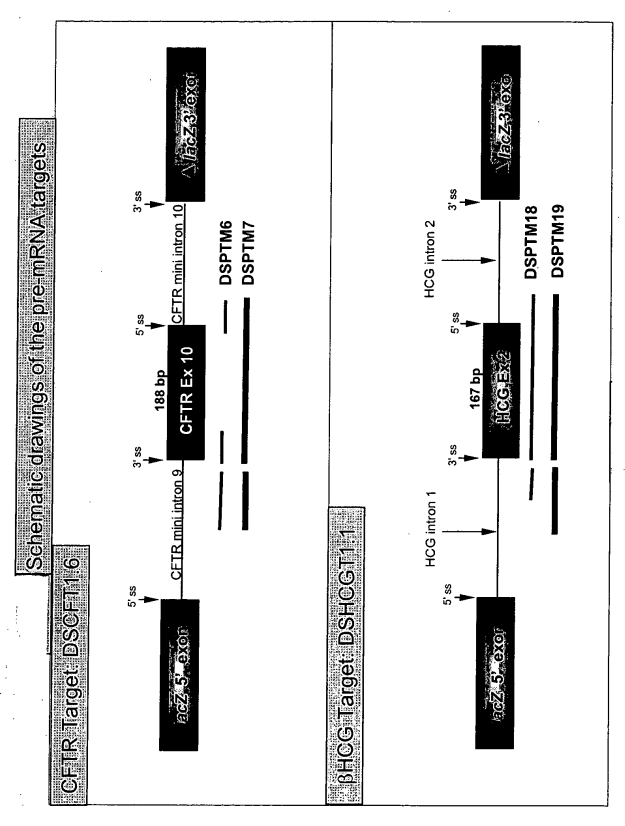
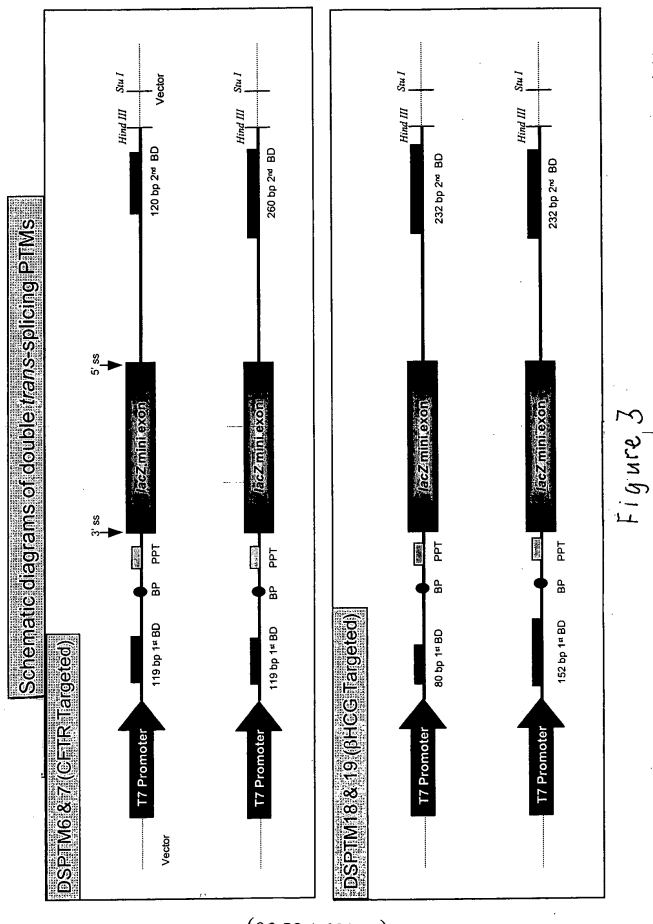
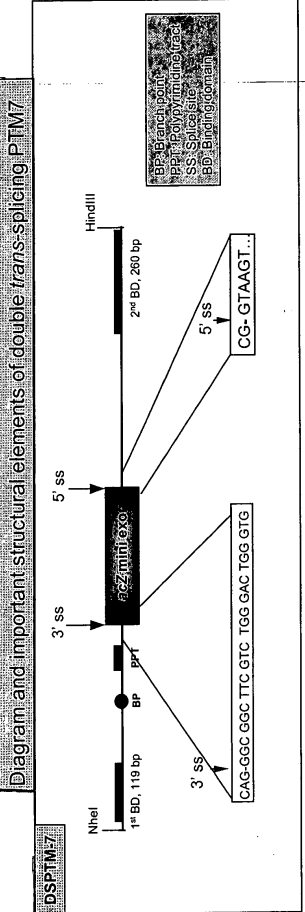


Figure 2

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A34759 (Sheet 4 of 10)



1st BD (119 bp): GATTCACTTGCTCCAATTATCATCCTAAGCAGAAGTGTATATTCTTATTTGTAAAGATTCTATTAACTCATTTGATT¢AAAATA TTTAAAATACTTCCTGTTTCATACTCTGCTATGCAC

Spac r sequences: AACATTATTATAACGTTGCTCGAA

BP, PPT and acceptor splice site: TACTAAC T GGTACC TCTTTTTTTTT GATATC CTGCAG GGC GGC TTC GTC TGG GAC TGG lacZ mini exon PPT

3, ss

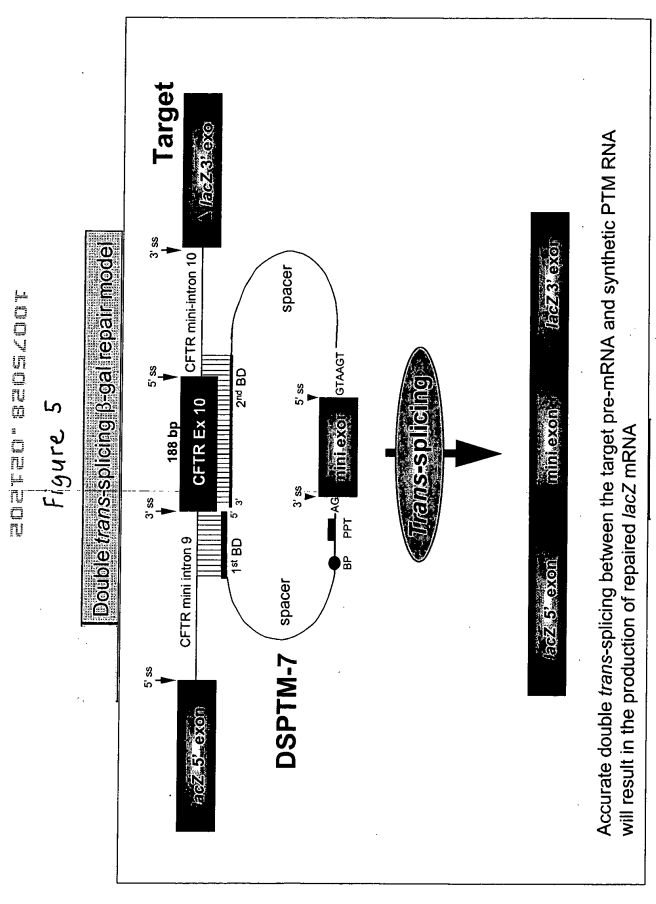
lacZ mini exon 5' ss

5' donor site and 2nd spacer sequence: IGA ACG GTAAGT GTTATCACCGATATGTGTCTAACCTGATTCGGGCCTTCGATACGCTAA GATCCACCGG

2nd BD (260 bp): TCAAAAAGTTTTCACATAATTTCTTACCTCTTCTTGAATTCATGCTTTGATGACGCTTCTGTATCTATATTCATCGAAA AAAAACCCTCTGAA77CTCCATTTCTCCCATAATCATCATTACAACTGAACTCTGGAAATAAAACCCATCATTATTAACTCA ACACCAATGATTTTTCTTTAATGGTGCCTGGCATAATCCTGGAAAACTGATAACACAATGAAATTCTTCCACTGTGCTTAA TTATCAAATCACGC

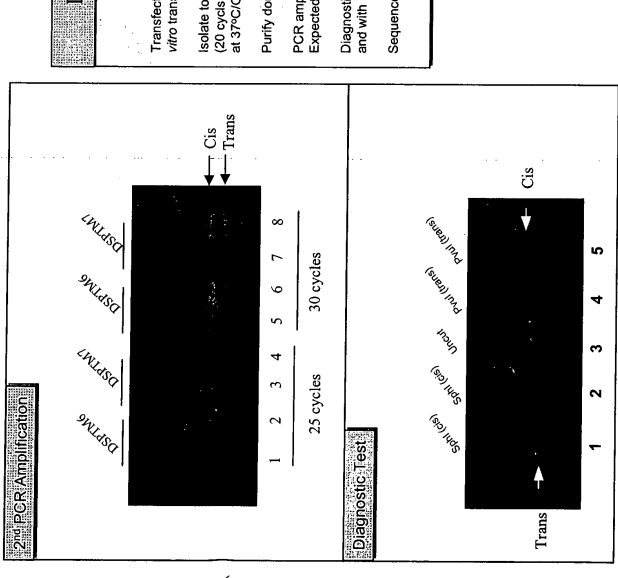
A34759 (Sheet 5 of 10)

Figure 4



A34759 (01 10)

Proof-of-principle of SMaRT using synthetic double splicing PTM RNA in 293T cells



DSPTM6 and 7 (CHTR targeted)

Methods

Transfect 293T cells with DSPTM6 and DSPTM7 in vitro transcribed, gel purified RNA (2.5-5.0 μg)

Isolate total RNA, cDNA synthesis (Lac6R), PCR amplification (20 cycls, KI-1F + Lac6R), digest with Sph I + Dde I (cis-specific) at 37°C/ON

Purify double trans-spliced product using Biotin-Lac21R probe

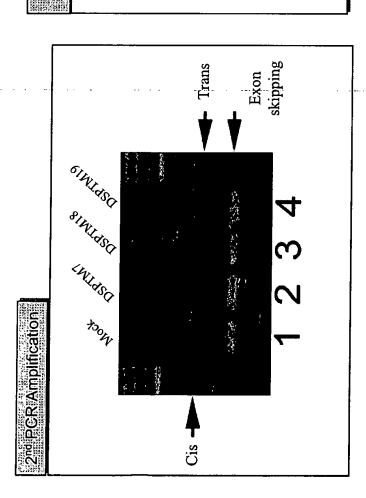
PCR amplify the captured trans-spliced product (KI-2F+Lac6R). Expected products: cis- 260bp; trans- 220 bp.

Diagnostic test: Digest PCR product with Pvu I (trans-specific) and with Sph I (cis-specific) at 37°C for 2-3 hr

Sequence to confirm the accuracy of double trans-splicing

Figure 6A

Proof-of-principle of SMaR√Tusing synthetic double splicing PTM RNA in stable cells



(Sheet 8 of 10)

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DSPTIM18 and 119 (HCG targeted)

Methods

Transfect DSHCGT1.1 stable cells with DSPTM7, DSPTM18 and DSPTM19 in vitro transcribed, gel purified RNA (2:5-5.0 µg)

Isolate total RNA, cDNA synthesis (Lac6R), PCR amplification (20 cycls, KI-1F + Lac6R), digest with Sph I + Dde I (cis-specific) at 37°C/ON

Purify double trans-spliced product using Biotin-Lac21R probe

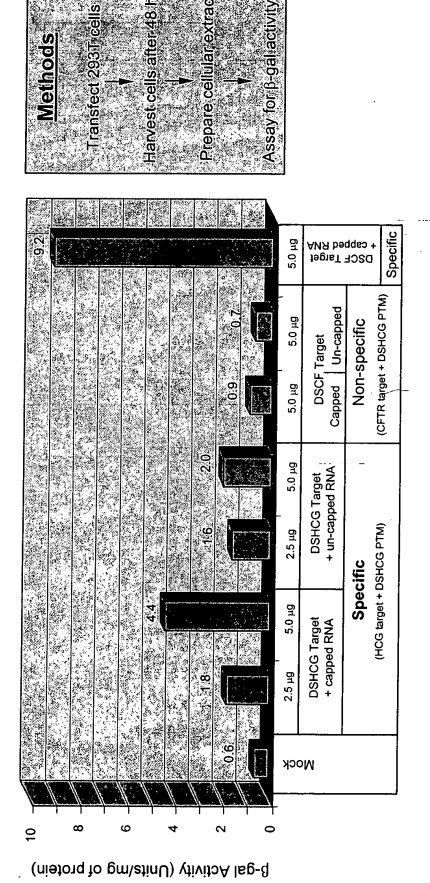
PCR amplify the captured *trans*-spliced product (KI-2F + Lac6R). Expected products: *cis*- 260bp; *trans*- 220 bp

Sequence to confirm the accuracy of double trans-splicing

Figure 6B

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Restoration of β-gal function through RNA transfection in 293T cells (Proof-of-concept for SMaRT RNA Therapeutics) Synthetic RNA; Double trans-splicing



(Sheet 10 of 10) 65745A

Figure 7